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(54) Title: A PROCESS AND SYSTEMS FOR THE EPOXIDATION OF AN OLEFIN

(57) Abstract: A process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a silver-based catalyst with the reaction modifier being present in a relative quantity Q which is the ratio of an effective molar quantity of active species of the reaction modifier present in the feed to an effective molar quantity of hydrocarbons present in the feed, and which process comprises the steps of:- operating at a first operating phase wherein the value of Q is Q1, and-subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase, such that the value of Q is Q2, whereby the value of the quotient Q2/Q1 is in the range of from 0.5 to 1.5; a system suitable for performing the process; and a computer program product and a computer system suitable for use in connection with the process.

A PROCESS AND SYSTEMS FOR THE EPOXIDATION OF AN OLEFIN

Field of the Invention

The invention relates to a process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a highly selective silver-based catalyst. The invention also relates to systems suitable for use in connection with the process.

Background of the Invention

Dased catalysts has been known for a long time. Conventional silver-based catalysts have provided the olefin oxides notoriously in a low selectivity. For example, when using conventional catalysts in the epoxidation of ethylene, the selectivity towards ethylene oxide, expressed as a fraction of the ethylene converted, does not reach values above the 6/7 or 85.7 mole-% limit. Therefore, this limit has long been considered to be the theoretically maximal selectivity of this reaction, based on the stoichiometry of the reaction equation

20 7 $C_2H_4 + 6 O_2 => 6 C_2H_4O + 2 CO_2 + 2 H_2O_7$

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cf. Kirk-Othmer's Encyclopedia of Chemical Technology, 3rd ed., Vol. 9, 1980, p. 445.

Modern silver-based catalysts however are more selective towards olefin oxide production. When using the modern catalysts in the epoxidation of ethylene the selectivity towards ethylene oxide can reach values above the 6/7 or 85.7 mole-% limit referred to. Such highly

selective catalysts, which may comprise as their active components silver, rhenium, at least one further element and optionally a rhenium co-promoter, are disclosed in EP-A-266015 and in several subsequent patent publications.

Besides better catalysts, reaction modifiers have been found which may be added to the feed to improve the selectivity (cf. for example EP-A-352850). Such reaction modifiers suppress the undesirable oxidation of olefin or olefin oxide to carbon dioxide and water, relative to the desired formation of olefin oxide, by a so-far unexplained mechanism. Suitable reaction modifiers are for example organic halides.

When applying a reaction modifier, the concentration of the reaction modifier in the feed may be chosen such that the selectivity is at the optimum value. The concentration at which the selectivity is at optimum may be found during the operation of the epoxidation process by a trial-and-error procedure, viz. by stepwise changing the reaction modifier supply rate and monitoring the effect on the selectivity. Such a procedure, however, would be cumbersome, and it would keep the process operating for some time at conditions which are less than the most economical. Moreover, the trial-and-error procedure would need to be redone when the feed composition has changed, in order to adjust the concentration of the reaction modifier to the new reaction conditions.

Summary of the Invention

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The present invention enables the operator of an epoxidation process to avoid unwanted variations in the selectivity when the feed composition changes by changing the concentration of the reaction modifier such that a relative quantity Q of the reaction modifier is maintained substantially, preferably completely, at a constant level. Herein, the relative quantity Q is basically the ratio of

the molar quantity of the reaction modifier in the feed to the molar quantity of hydrocarbons in the feed. words, the teaching of the present invention is that the concentration of the reaction modifier needed to achieve a certain effect on the selectivity is proportional to the concentration of the hydrocarbons present in the feed and, as a consequence, this effect on the selectivity may be preserved when changing the feed composition by also changing the reaction modifier concentration proportionally or substantially proportionally with any change in the hydrocarbon concentration. This is independent of any change in the feed composition other than changes relating to the hydrocarbons and/or the reaction modifier. Thus, it is a benefit of the present invention that it allows the epoxidation process to be controlled significantly more simply and more smoothly than without the invention.

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It has also been found that in a quantitative sense there may be differences in the behavior of the various hydrocarbons which may be present in the reaction mixture, and it is therefore more preferred, when calculating Q, to replace the molar quantity of hydrocarbons by a -so-called-effective molar quantity of hydrocarbons. The effective molar quantity of hydrocarbons in the feed can be calculated from the feed composition, as set out hereinafter, such that it accounts for differences in the behavior between the various hydrocarbons present.

Further, it has been found that in a quantitative sense there may also be differences in the behavior of different reaction modifiers, while in practice a mixture of reaction modifiers is frequently present. Therefore it may be preferred, when calculating Q, also to replace the molar quantity of the reaction modifier by a -so-called- effective molar quantity of active species of the reaction modifier. The effective molar quantity of active species of the reaction modifier in the feed can be calculated from the

feed composition, as set out hereinafter, such that it accounts for the differences in the behavior of different reaction modifiers.

Not wishing to be bound by theory, it is thought that, 5 unlike other components of the feed, the hydrocarbons (for example, the olefin and saturated hydrocarbons, if present) have an ability to remove or strip reaction modifier from the catalyst. It is the teaching of this invention that, in order to maintain the effects of the reaction modifier, it is the concentration of the modifier's active species on the catalyst which needs to be maintained, as opposed to the concentration of the reaction modifier at places in the reaction mixture other than the catalyst surface. Differences in the ability of the various hydrocarbons in the removing/stripping process and differences in the effectiveness of the various reaction modifiers and their susceptibility to the removing/stripping process are thought to be accounted for by calculating and applying the effective molar quantities, as explained hereinbefore.

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20 Accordingly, the present invention provides a process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a silver-based catalyst with the reaction modifier being present in a relative quantity Q 25 which is the ratio of an effective molar quantity of active species of the reaction modifier present in the feed to an effective molar quantity of hydrocarbons present in the feed, and which process comprises the steps of:

- operating at a first operating phase wherein the value of Q is Q_1 , and
 - subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase, such that the value of Q is Q_2 , whereby the value of the quotient Q_2/Q_1 is in the range of from 0.5 to 1.5.

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In a particular embodiment, the second phase is operated at a hydrocarbon composition and a reaction modifier composition of the feed of which at least one is different from the hydrocarbon composition and the reaction modifier composition of the feed employed in the first operating phase.

The invention also provides a system suitable for performing the process of this invention, which system comprises a reactor which holds a silver-based catalyst, means for feeding to the reactor a feed comprising the olefin, oxygen and a reaction modifier, and feed control means for controlling the feed and/or the feed composition, comprising modifier control means for controlling the reaction modifier being present in the feed in a relative quantity Q which is the ratio of an effective molar quantity of active species of the reaction modifier present in the feed to an effective molar quantity of hydrocarbons present in the feed, and which feed control means are configured such as to control the process steps of:

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- operating at a first operating phase wherein the value of Q is Q_1 , and
- subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase, such that the value of Q is Q_2 , whereby the value of the quotient Q_2/Q_1 is in the range of from 0.5 to 1.5.

The invention also provides a computer program product which comprises a computer readable medium and a computer readable program code, recorded on the computer readable medium, suitable for instructing a data processing system of a computer system to execute calculations for the process of this invention.

The invention also provides a computer system which comprises the computer program product of this invention and a data processing system configured to receive instructions read from the computer program product.

The invention also provides, in more general terms, a process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a silver-based catalyst, which process comprises the steps of:

10 - operating at a first operating phase, and

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- subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase, such that the concentration of active species of the reaction modifier on the catalyst is substantially unchanged.

Detailed Description of the Invention

Although the present epoxidation process may be carried out in many ways, it is preferred to carry it out as a gas phase process, i.e. a process in which the feed is contacted in the gas phase with the catalyst which is present as a solid material, typically in a packed bed. Generally the process is carried out as a continuous process. Frequently, in commercial scale operation, the process of the invention may involve a quantity of catalyst which is at least 10 kg, for example at least 20 kg, frequently in the range of from 10^2 to 10^7 kg, more frequently in the range of from 10^6 kg.

The olefin for use in the present epoxidation process may be any olefin, such as an aromatic olefin, for example styrene, or a di-olefin, whether conjugated or not, for example 1,9-decadiene or 1,3-butadiene. Typically, the olefin is a monoolefin, for example 2-butene or isobutene. Preferably, the olefin is a mono- α -olefin, for example 1-butene or propylene. The most preferred olefin is ethylene.

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The olefin concentration in the feed is not material to this invention and may be selected within a wide range.

Typically, the olefin concentration in the feed will be at most 80 mole-%, relative to the total feed. Preferably, it will be in the range of from 0.5 to 70 mole-%, in particular from 1 to 60 mole-%, on the same basis. As used herein, the feed is considered to be the composition which is contacted with the catalyst.

The present epoxidation process may be air-based or oxygen-based, see Kirk-Othmer's Encyclopedia of Chemical Technology, 3rd ed., Vol. 9, 1980, p. 445-447. In the air-based process air or air enriched with oxygen is employed as the source of the oxidizing agent while in the oxygen-based processes high-purity (>95 mole-%) oxygen is employed as the source of the oxidizing agent. Presently most epoxidation plants are oxygen-based and this is a preferred embodiment of the present invention.

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The oxygen concentration in the feed is not material to this invention and may be selected within a wide range. However, in practice, oxygen is generally applied at a concentration which avoids the flammable regime. Typically, the concentration of oxygen applied will be within the range of from 1 to 15 mole-%, more typically from 2 to 12 mole-% of the total feed.

In order to remain outside the flammable regime, the concentration of oxygen in the feed may be lowered as the concentration of the olefin is increased. The actual safe operating ranges depend, along with the feed composition, also on the reaction conditions such as the reaction temperature and the pressure.

The reaction modifier is present in the feed for increasing the selectivity, suppressing the undesirable oxidation of olefin or olefin oxide to carbon dioxide and water, relative to the desired formation of olefin oxide.

Many organic compounds, especially organic halides and organic nitrogen compounds, may be employed as the reaction modifier. Nitrogen oxides, hydrazine, hydroxylamine or ammonia may be employed as well. It is frequently considered that under the operating conditions of olefin epoxidation the nitrogen containing reaction modifiers are precursors of nitrates or nitrites, i.e. they are so-called nitrate- or nitrite-forming compounds (cf. e.g. EP-A-3642, US-A-4822900).

Organic halides are the preferred reaction modifiers, in particular organic bromides, and more in particular organic chlorides. Preferred organic halides are chlorohydrocarbons or bromohydrocarbons. More preferably they are selected from the group of methyl chloride, ethyl chloride, ethylene dichloride, ethylene dibromide, vinyl chloride or a mixture thereof. Most preferred reaction modifiers are ethyl chloride and ethylene dichloride.

Suitable nitrogen oxides are of the general formula NO_x wherein x, which denotes the ratio of the number of oxygen atoms to the number of nitrogen atoms, is in the range of from 1 to 2. These nitrogen oxides include for example NO, N_2O_3 and N_2O_4 . Suitable organic nitrogen compounds are nitro compounds, nitroso compounds, amines, nitrates and nitrites, for example nitromethane, 1-nitropropane or 2-nitropropane. In preferred embodiments, nitrate- or nitrite-forming compounds, e.g. nitrogen oxides and/or organic nitrogen compounds, are used together with an organic halide, in particular an organic chloride.

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Although the reaction modifier may be supplied as a single compound, upon contact with the catalyst a variety of compounds may be formed which function as reaction modifier, and which may be present in the feed if a recycle is applied. For example, when applying ethyl chloride in an ethylene oxide process, the feed may in practice comprise

ethyl chloride, vinyl chloride, ethylene dichloride and methyl chloride.

The reaction modifiers are generally effective when used in low concentration in the feed, for example up to 5 0.1 mole-%, relative to the total feed, for example from 0.01×10^{-4} to 0.01 mole-%. In particular when the olefin is ethylene, it is preferred that the reaction modifier is present in the feed at a concentration of from 0.05×10^{-4} to 50×10^{-4} mole-%, in particular from 0.2×10^{-4} to 30×10^{-4} mole-%, relative to the total feed.

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In addition to the olefin, oxygen and the reaction modifier, the feed may contain one or more optional components, such as carbon dioxide, water, inert gases and saturated hydrocarbons. Carbon dioxide and water are byproducts of the epoxidation process. Carbon dioxide generally has an adverse effect on the catalyst activity. Typically, a concentration of carbon dioxide in the feed in excess of 25 mole-%, preferably in excess of 10 mole-%, relative to the total feed, is avoided. A concentration of carbon dioxide as low as 1 mole-% or lower, relative to the total feed, may be employed. Water may be introduced in the feed as a result of the recovery of olefin oxide and carbon dioxide from the reaction product. Water generally has an adverse effect on the catalyst activity. Typically, a concentration of water in the feed in excess of 3 mole-%, preferably in excess of 1 mole-%, relative to the total feed, is avoided. A concentration of water as low as 0.2 mole-% or lower, relative to the total feed, may be employed. Inert gas, for example nitrogen or argon, or a mixture thereof, may be present in the feed in a concentration of from 0.5 to 95 mole-%. In an air based process inert gas may be present in the feed in a concentration of from 30 to 90 mole-%, typically from 40 to 80 mole-%. In an oxygen based process inert gas may be

present in the feed in a concentration of from 0.5 to 30 mole-%, typically from 1 to 15 mole-%. Suitable saturated hydrocarbons are propane and cyclopropane, and in particular methane and ethane. If saturated hydrocarbons are present, 5 they may be present in a quantity of up to 80 mole-%, relative to the total feed, in particular up to 75 mole-%. Frequently they are present in a quantity of at least 30 mole-%, more frequently at least 40 mole-%. Saturated hydrocarbons may be added to the feed in order to increase the oxygen flammability limit.

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The relative quantity Q of the reaction modifier is the ratio of the effective molar quantity of active species of the reaction modifier present in the feed to the effective molar quantity of hydrocarbons present in the feed, both molar quantities being expressed in the same units, for example as mole-%, based on the total feed.

When the reaction modifier is a halogen compound, for the purpose of calculating the effective molar quantity of active species of the reaction modifier and the value of Q, the number of active species is deemed to be the number of halogen atoms, and when the reaction modifier is a nitrateor nitrite-forming compound, the number of active species is deemed to be the number of nitrogen atoms. This implies, for example, that 1 mole of ethylene dichloride provides 2 moles of active species, i.e. all of the chlorine atoms present provide an active species. On the other hand, it has also been found that reaction modifiers which are methyl compounds, such as methyl chloride and methyl bromide, are less responsive and therefore from 2 to 5 moles, in particular from 2.5 to 3.5 moles, suitably 3 moles of the methyl compounds may be deemed to provide 1 mole of the active species. This number may be determined and verified by routine experimentation, and -without wishing to be bound by theory- it is believed that this number is higher as the methyl compound in question has a lesser ability to split

off the heteroatom in question (for example the halogen or nitrogen atom). Thus, for example, when the feed comprises 2×10^{-4} mole-% of ethyl chloride, 3×10^{-4} mole-% of vinyl chloride, 1×10^{-4} mole-% of ethylene dichloride and 1.5×10^{-4} mole-% of methyl chloride, the effective molar quantity of active species of the reaction modifier may be calculated to amount to $2\times10^{-4}\times1+3\times10^{-4}\times1+1\times10^{-4}\times2+1.5\times10^{-4}\times1/_3=7.5\times10^{-4}$ mole-%.

Summarizing, the effective molar quantity of active species of the reaction modifier present in the feed may be calculated by multiplying the molar quantity of each of the reaction modifiers present in the feed with a factor, and adding up the resulting multiplication products, wherein each factor represents the number of active heteroatoms, in particular halogen atoms and/or nitrogen atoms, present per molecule of the reaction modifier in question, on the understanding that the factor for a reaction modifier which is a methyl compound may be in the range of from $^{1}/_{5}$ to $^{1}/_{2}$, more typically from $^{1}/_{3.5}$ to $^{1}/_{2.5}$, suitably $^{1}/_{3}$.

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The hydrocarbons present in the feed comprise the olefin and any saturated hydrocarbon present. As indicated herein-before, it is thought that the hydrocarbons present in the feed have the ability to remove/strip reaction modifier from the catalyst surface and the extent to which they have this ability may differ for the various hydrocarbons. In order to account for these differences (relative to ethylene), the molar quantity of each of the hydrocarbons present is multiplied with a factor, before the molar quantities are added up to calculate the effective molar quantity of the hydrocarbons. Herein, the factor of ethylene is 1, by definition; the factor for methane may be at most 0.5, or at most 0.4, typically in the range of from 0 to 0.1; the factor for ethane may be in the range of from 50 to 150,

more typically from 70 to 120; and the factor for higher hydrocarbons (i.e. having at least 3 carbon atoms) may be in the range of from 10 to 10000, more typically from 50 to 2000. Such factors may be determined and verified by routine experimentation, and -without wishing to be bound by theoryit is believed that the factor is higher as the hydrocarbon in question has a greater ability to form radicals. Suitable factors for methane, ethane, propane and cyclopropane, relative to ethylene, are 0.3, 85, 1000, and 60, respectively. As an example, when the feed comprises 30 mole-% ethylene, 40 mole-% of methane, 0.4 mole-% of ethane and 0.0001 mole-% of propane, the effective molar quantity of the hydrocarbons may be calculated to amount to $30 \times 1 + 40 \times 0.1 + 0.4 \times 85 + 0.0001 \times 1000 = 68.1 \text{ mole-}\%$.

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It is noted that when ethylene oxide is produced from ethylene without further hydrocarbons being present, the effective molar quantity of the hydrocarbons equals the actual molar quantity, and that the addition of ethane or higher hydrocarbons to an ethylene feed contributes

20 significantly to the effective molar quantity, whereas there is relatively little contribution from any methane added.

Eligible values of Q are at least 1×10^{-6} , and in particular at least 2×10^{-6} . Eligible values of Q are at most 100×10^{-6} , and in particular at most 50×10^{-6} .

In any operating phase of the process the concentration of the reaction modifier and, thus, the value of Q may be adjusted so as to achieve an optimal selectivity towards the olefin oxide formation, for the prevailing reaction conditions. This relates in particular to embodiments of this invention in which the catalyst is a highly selective silver based catalyst, as defined hereinafter.

In accordance with this invention, when the feed composition changes, the concentration of the reaction

modifier may be changed as well, such that the value of Q is not substantially changed, which means that Q_1 is approximately equal, or preferably equal to Q_2 . Preferably, the value of the quotient Q_2/Q_1 is in the range of from 0.8 to 1.2, in particular from 0.9 to 1.1, more in particular from 0.95 to 1.05. Most preferably the value of the quotient Q_2/Q_1 equals 1.

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In this way the present invention enables to determine by calculation a desirable composition of the reaction modifier(s) or the hydrocarbon(s) in the feed which may be applied in the second operating phase of the epoxidation process, in response to a change in the feed composition as regards the quantity or type of the components of the feed. In a preferred embodiment, a desirable concentration of the reaction modifier(s) in the feed is calculated which may be applied in the second operating phase, in response to a change in the quantity or type of hydrocarbon(s) present in the feed. In another embodiment, a desirable concentration of the hydrocarbon(s) in the feed is calculated which may be applied in the second operating phase, in response to a change in the quantity or type of the reaction modifier(s) present in the feed. In again another embodiment, a desirable concentration of the reaction modifier(s) in the feed is calculated which may be applied in the second operating phase, in response to a change in the type of the reaction modifier(s) present in the feed, while it is preferred not to change the quantity or type of the hydrocarbon(s). In again another embodiment, a desirable concentration of the hydrocarbon(s) in the feed is calculated which may be applied in the second operating phase, in response to a change in the type of hydrocarbon(s) present in the feed, while it is preferred not to change the quantity or type of the reaction modifier(s).

There may be various reasons for the occurrence of changes in the feed composition. For example, by changing

the oxygen concentration or the olefin concentration the production rate of the olefin oxide may be controlled, or changing the concentration of saturated hydrocarbons may change the flammability limits. The carbon dioxide 5 concentration in the feed may change as a result of catalyst ageing. The concentration of inert gases in the feed may change as a result of a change in the concentration of such gases in the oxygen supplied to the process. Further, when a high selectivity catalyst is applied, the selectivity may be improved at an advanced stage of ageing of the catalyst by increasing the ethylene content of the feed (cf. US-6372925-B1 and WO-A-01/96324, i.e. non-prepublished PCT patent application PCT/US01/18097). By applying this invention when the process is operating at optimal selectivity, the change in the feed composition leads to less deviation of the selectivity from the optimum or even the selectivity may be maintained at the optimum level. By applying this invention when the process is not operating at optimal selectivity, application of this invention prevents the process from deviating further away from the optimum selectivity.

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Any change in the feed composition may be gradual, or step wise, and any change in the feed composition may be accompanied with a corresponding change in the concentration of the reaction modifier, such that the value of Q is substantially unchanged or it is kept constant. Generally, the change in the feed composition is accompanied with a cocurrent change in the concentration of the reaction modifier.

The present epoxidation process may be carried out using reaction temperatures selected from a wide range. Preferably the reaction temperature is in the range of from 180 to 340 °C, more preferably in the range of from 190 to 325 °C, in particular in the range of from 200 to 300 °C. Preferably, as the catalyst ages, the reaction temperature

is slowly increased as to compensate for a reduction in the activity of the catalyst. However, it is preferred that the change in the feed composition occurs substantially without a change in the reaction temperature. Frequently a change in reaction temperature, if any, accompanying the change in the feed composition, may be less than 10 °C, more frequently less than 5 °C, in particular less than 2 °C. Most preferably, the change in the feed composition is effected without a change in the reaction temperature at all.

Generally, the silver based catalyst is a supported catalyst. The support may be selected from a wide range of inert support materials. Such support materials may be natural or artificial inorganic materials and they include silicon carbide, clays, pumice, zeolites, charcoal and alkaline earth metal carbonates, such as calcium carbonate. Preferred are refractory support materials, such as alumina, magnesia, zirconia and silica. The most preferred support material is α-alumina.

The support material is preferably porous and has

20 preferably a surface area, as measured by the B.E.T. method, of less than 20 m²/g and in particular from 0.05 to 20 m²/g. More preferably the B.E.T. surface area of the support is in the range of 0.1 to 10, in particular from 0.1 to 3.0 m²/g. As used herein, the B.E.T. surface area is deemed to have been measured by the method as described in Brunauer, Emmet and Teller in J. Am. Chem. Soc. 60 (1938) 309-316.

The silver based catalyst preferably comprises silver and a further element or compound thereof. Eligible further elements are selected from the group of nitrogen, sulfur, phosphorus, boron, fluorine, Group IA metals, Group IIA metals, rhenium, molybdenum, tungsten, chromium, titanium, hafnium, zirconium, vanadium, thallium, thorium, tantalum, niobium, gallium and germanium and mixtures thereof.

Preferably the Group IA metals are selected from lithium,

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potassium, rubidium and cesium. Most preferably the Group IA metal is lithium, potassium and/or cesium. Preferably the Group IIA metals are selected from calcium and barium. Where possible, the further element may suitably be provided as an oxyanion, for example as a sulfate, borate, perrhenate, molybdate or nitrate, in salt or acid form.

It is preferred to employ highly selective silver-based catalysts. The highly selective silver-based catalysts comprise, in addition to silver, one or more of rhenium, molybdenum, tungsten, a Group IA metal, and a nitrate- or nitrite-forming compound, which may each be present in a quantity of from 0.01 to 500 mmole/kg, calculated as the element (rhenium, molybdenum, tungsten, the Group IA metal or nitrogen) on the total catalyst. The nitrate- or nitrite-forming compounds and particular selections of nitrate- or nitrite-forming compounds are as defined hereinbefore. Rhenium, molybdenum, tungsten or the nitrate- or nitrite-forming compound may suitably be provided as an oxyanion, i.e. as a perrhenate, molybdate, tungstate or nitrate, in salt or acid form.

Of special preference are the highly selective silver based catalysts which comprise rhenium in addition to silver. Such catalysts are known from EP-A-266015. Broadly, they comprise silver, rhenium or compound thereof, the further element (as defined hereinbefore) other than rhenium or compound thereof and optionally a rhenium co-promoter which may be selected from one or more of sulfur, phosphorus, boron, and compounds thereof.

Preferred amounts of the components of the silver based 30 catalysts are, when calculated as the element on the total catalyst:

- silver from 10 to 500 g/kg,

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- if present, rhenium from 0.01 to 50 mmole/kg,
- if present, the further element or elements from 0.1 to

500 mmole/kg each, and,

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- if present, the rhenium co-promoter or co-promoters from 0.1 to 30 mmole/kg each.

The present epoxidation process is preferably carried out at a reactor inlet pressure in the range of from 1000 to 4000 kPa. "GHSV" or Gas Hourly Space Velocity is the unit volume of gas at normal temperature and pressure (0 °C, 1 atm, i.e. 101.3 kPa) passing over one unit volume of packed catalyst per hour. Preferably, when the epoxidation 10 process is as a gas phase process involving a packed catalyst bed, the GHSV is in the range of from 1500 to 10000 Nl/(l.h). Preferably, the process of this invention is carried out at a work rate in the range of from 0.5 to 10 kmole olefin oxide produced per m³ of catalyst per hour, in particular 0.7 to 8 kmole olefin oxide produced per m³ of 15 catalyst per hour, for example 5 kmole olefin oxide produced per m³ of catalyst per hour. As used herein, the work rate is the amount of the olefin oxide produced per unit volume of catalyst per hour and the selectivity is the molar quantity of the olefin oxide formed relative to the molar quantity of 20 the olefin converted.

The olefin oxide produced may be recovered from the reaction product by using methods known in the art, for example by absorbing the olefin oxide from a reactor outlet stream in water and optionally recovering the olefin oxide from the aqueous solution by distillation. At least a portion of the aqueous solution containing the olefin oxide may be applied in a subsequent process for converting the olefin oxide into a 1,2-diol or a 1,2-diol ether.

The olefin oxide produced in the present epoxidation process may be converted into a 1,2-diol or into a 1,2-diol ether. As this invention leads to a more attractive process for the production of the olefin oxide, it concurrently leads to a more attractive process which comprises producing

the olefin oxide in accordance with the invention and the subsequent use of the obtained olefin oxide in the manufacture of the 1,2-diol and/or 1,2-diol ether.

The conversion into the 1,2-diol or the 1,2-diol ether may comprise, for example, reacting the olefin oxide with water, suitably using an acidic or a basic catalyst. For example, for making predominantly the 1,2-diol and less 1,2diol ether, the olefin oxide may be reacted with a ten fold molar excess of water, in a liquid phase reaction in presence of an acid catalyst, e.g. 0.5-1.0 %w sulfuric acid, 10 based on the total reaction mixture, at 50-70 °C at 1 bar absolute, or in a gas phase reaction at 130-240 °C and 20-40 bar absolute, preferably in the absence of a catalyst. If the proportion of water is lowered the proportion of 1,2diol ethers in the reaction mixture is increased. The 1,2diol ethers thus produced may be a di-ether, tri-ether, tetra-ether or a subsequent ether. Alternative 1,2-diol ethers may be prepared by converting the olefin oxide with an alcohol, in particular a primary alcohol, such as 20 methanol or ethanol, by replacing at least a portion of the water by the alcohol.

The 1,2-diol and the 1,2-diol ether may be used in a large variety of industrial applications, for example in the fields of food, beverages, tobacco, cosmetics, thermoplastic polymers, curable resin systems, detergents, heat transfer systems, etc.

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Unless specified otherwise, the organic compounds mentioned herein, for example the olefins, 1,2-diols, 1,2-diol ethers and reaction modifiers, have typically at most 40 carbon atoms, more typically at most 20 carbon atoms, in particular at most 10 carbon atoms, more in particular at most 6 carbon atoms. As defined herein, ranges for numbers of carbon atoms (i.e. carbon number) include the numbers specified for the limits of the ranges.

The computer program product of this invention comprises a computer readable medium and a computer readable program code, recorded on the computer readable medium, suitable for instructing a data processing system to execute the calculations in connection with the process of this invention. In a preferred embodiment, the computer program product comprises, in addition, a computer readable program code, recorded on the computer readable medium, suitable for instructing the data processing system to control the process of this invention. The computer readable medium may be readable, for example, by means of an optical system or by means of a magnetic system. The computer program product may be in the form of a disk which is a permanent entity of the computer system of this invention, or it may be a disk which is insertable into the computer system. control means may be configured such that they communicate with the computer system of this invention facilitating the control of the process steps of the process of this invention.

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A catalyst, as defined in EP-A-266015, containing silver, rhenium, cesium, lithium and sulfur on α -alumina and having a theoretical selectivity S_0 of 93% in the fresh state is employed in the following experiment. The above value of S_0 was determined by measuring selectivities in a range of gas hourly space velocities, each time at 30% ethylene, 8% oxygen, 5% carbon dioxide and 1400 kPa, the reaction temperature being 260 °C, and extrapolating back to zero oxygen conversion.

In the experiment ethylene oxide is produced as follows. A 1-kg sample of the catalyst is loaded into a tubular reactor consisting of a stainless steel tube. The tube is immersed in a boiling kerosene containing cooling

jacket and the ends are connected to a gas flow system. The inlet gas flow rate is adjusted as to achieve a gas hourly space velocity of 6800 Nl /(l.h). The inlet pressure is 2100 kPa (absolute). The feed to the reactor comprises ethylene at a concentration of 28 mole-%, oxygen at a concentration of 8 mole-%, carbon dioxide at a concentration of 3 mole-%, ethane at a concentration of 0.5 mole-%, ethyl chloride at a concentration of 3×10^{-4} mole-%, ethylene dichloride at a concentration of 0.5×10^{-4} mole-%, vinyl chloride at a concentration of 1×10^{-4} mole-%, and methyl chloride at a concentration of 2×10^{-4} mole-%, and the remainder of the feed is nitrogen. The reaction temperature is 250 °C. The selectivity to ethylene oxide is optimal in respect of the chlorides. The value of Q (i.e. Q_1) equals $(3\times10^{-4}\times1$ + $0.5 \times 10^{-4} \times 2 + 1 \times 10^{-4} \times 1 +$ $2\times10^{-4} \times {}^{1}/_{3}$) / (28 × 1 + 0.5 × 85) = 8.04×10⁻⁶.

At a certain point in time the feed composition changes such that ethylene is present at a concentration of 25 mole-% and ethane at a concentration of 0.7 mole-%, the concentration of the other constituents being held constant, except ethyl chloride. In order to keep the selectivity at an optimum level, the ethyl chloride concentration is adjusted to 4.1×10^{-4} mole-%. After these changes, the value of Q (i.e. Q₂) equals $(4.1 \times 10^{-4} \times 1 + 0.5 \times 10^{-4} \times 2 + 1 \times 10^{-4} \times 1 + 2 \times 10^{-4} \times \frac{1}{3})$

 $(25 \times 1 + 0.7 \times 85) = 8.01 \times 10^{-6}$.

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CLAIMS

A process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a silver-based catalyst with the reaction modifier being present in a relative quantity Q which is the ratio of an effective molar quantity of active species of the reaction modifier present in the feed to an effective molar quantity of hydrocarbons present in the feed, and which process comprises the steps of:

- operating at a first operating phase wherein the value of Q is Q_1 , and
- subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase, such that the value of Q is Q_2 , whereby the value of the quotient Q_2/Q_1 is in the range of from 0.5 to 1.5.
- 20 2. A process as claimed in claim 1, wherein the olefin is ethylene, and wherein the reaction modifier comprises an organic chloride and optionally a nitrate- or nitrite-forming compound.
- 3. A process as claimed in claim 2, wherein the reaction modifier consists of chlorohydrocarbons having up to 10 carbon atoms, in particular up to 6 carbon atoms, which comprise one or more of methyl chloride, ethyl chloride, ethylene dichloride and vinyl chloride.
- 4. A process as claimed in any of claims 1-3, wherein
 30 the effective molar quantity of active species of the
 reaction modifier present in the feed can be calculated by
 multiplying the molar quantity of each of the reaction
 modifiers present in the feed with a factor, and adding up
 the resulting multiplication products, wherein each factor

represents the number of active heteroatoms, in particular sulfur and/or nitrogen atoms, present per molecule of the reaction modifier in question, on the understanding that the factor for a reaction modifier which is a methyl compound, if present, is in the range of from $^{1}/_{5}$ to $^{1}/_{2}$, in particular in the range of from $^{1}/_{3.5}$ to $^{1}/_{2.5}$.

5. A process as claimed in any of claims 1-4, wherein the effective molar quantity of hydrocarbons present in the feed can be calculated by multiplying the molar quantity of each of the hydrocarbons present in the feed with a factor, and adding up the resulting multiplication products, wherein the factor for methane, if present, is in the range of from 0.1 to 0.5; the factor for ethane, if present, is in the range of from 50 to 150; and the factor for hydrocarbons having at least 3 carbon atoms, if present, is in the range of from 10 to 10000, all factors being relative to the factor for ethylene, which equals 1, by definition.

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- 6. A process as claimed in claim 5, wherein the factor for methane, if present, is in the range of from 0 to 0.4; the factor for ethane, if present, is in the range of from 70 to 120; and the factor for the hydrocarbons having at least 3 carbon atoms, if present, is in the range of from 50 to 2000, all factors being relative to the factor for ethylene, which equals 1, by definition.
- 25 7. A process as claimed in any of claims 1-6, wherein the relative quantity Q is in the range of from 1×10^{-6} to 100×10^{-6} , in particular in the range of from 2×10^{-6} to 50×10^{-6} .
 - 8. A process as claimed in any of claims 1-7, wherein the hydrocarbons present in the feed comprise one or more of methane, ethane, propane and cyclopropane, in addition to the olefin.
 - 9. A process as claimed in any of claims 1-8, wherein such a value of Q_1 is employed that the selectivity towards

the olefin oxide formation is optimal.

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10. A process according to any of claims 1-9, wherein the value of the quotient Q_2/Q_1 is in the range of from 0.8 to 1.2, in particular in the range of from 0.9 to 1.1, more in particular in the range of from 0.95 to 1.05.

- 11. A process according to any of claims 1-10, wherein the second phase is operated at a hydrocarbon composition and a reaction modifier composition of the feed of which at least one is different from the hydrocarbon composition and the reaction modifier composition of the feed employed in the first operating phase.
- 12. A process according to claim 11, wherein the concentration of the reaction modifier(s) in the feed applied in the second operating phase is calculated, in response to a change in the quantity or type of hydrocarbon(s) present in the feed.
- A process according to any of claims 1-12, wherein the silver based catalyst comprises silver and a further element or compound thereof, which further element is
 selected from the group of nitrogen, sulfur, phosphorus, boron, fluorine, Group IA metals, Group IIA metals, rhenium, molybdenum, tungsten, chromium, titanium, hafnium, zirconium, vanadium, thallium, thorium, tantalum, niobium, gallium and germanium and mixtures thereof, on a support, in
 particular an α-alumina support.
 - 14. A process according to claim 13, wherein the silver based catalyst comprises silver, rhenium or compound thereof, a further element or compound thereof which further element is selected from the group of nitrogen, sulfur, phosphorus, boron, fluorine, Group IA metals, Group IIA metals, molybdenum, tungsten, chromium, titanium, hafnium, zirconium, vanadium, thallium, thorium, tantalum, niobium, gallium and germanium and mixtures thereof, and optionally a rhenium co-promoter which may be selected from one or more

of sulfur, phosphorus, boron, or compound thereof, on a support material, in particular an α -alumina support.

- 15. A process for the epoxidation of an olefin, which process comprises reacting a feed comprising the olefin, oxygen and a reaction modifier in the presence of a silver-based catalyst, which process comprises the steps of:
- operating at a first operating phase, and

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- subsequently operating at a second operating phase wherein the feed composition is different from the feed composition employed in the first operating phase such that the concentration of active species of the reaction modifier on the catalyst is substantially unchanged.
- 16. A method for making a 1,2-diol or a 1,2-diol ether comprising converting an olefin oxide into the 1,2-diol or the 1,2-diol ether, wherein the olefin oxide has been obtained by a process for the production of olefin oxide according to any of claims 1-15.
- A system suitable for performing a process 17. according to any of claims 1-15, which system comprises a reactor which holds a silver-based catalyst, means for 20 feeding to the reactor a feed comprising the olefin, oxygen and a reaction modifier, and feed control means for controlling the feed and/or the feed composition, comprising modifier control means for controlling the reaction modifier being present in the feed in a relative quantity Q which is 25 the ratio of an effective molar quantity of active species of the reaction modifier present in the feed to an effective molar quantity of hydrocarbons present in the feed, and which feed control means are configured such as to control 30 the process steps of:
 - operating at a first operating phase wherein the value of Q is Q₁, and
 - subsequently operating at a second operating phase
 wherein the feed composition is different from the feed

composition employed in the first operating phase, such that the value of Q is Q_2 , whereby the value of the quotient Q_2/Q_1 is in the range of from 0.5 to 1.5.

- 18. A computer program product which comprises a computer readable medium and a computer readable program code, recorded on the computer readable medium, suitable for instructing a data processing system of a computer system to execute calculations for the process according to any of claims 1-15.
- 10 19. A computer program product as claimed in claim 20, which comprises, in addition, a computer readable program code, recorded on the computer readable medium, suitable for instructing the data processing system to control a process according to any of claims 1-15.
- 15 20. A computer system which comprises a computer program product and a data processing system configured to receive instructions read from the computer program product, wherein the computer program product is as claimed in claim 18 or 19.

INTERNATIONAL SEARCH REPORT

Internation Pilication No PCT/US 02/37022

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07D301/10									
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According to International Patent Classification (IPC) or to both national classification and IPC									
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	cumentation searched (classification system followed by classification	n symbols)							
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Electronic data base consulted during the international search (name of data base and, where practical, search terms used)									
EPO-In	ternal, WPI Data, PAJ								
	ENTS CONSIDERED TO BE RELEVANT		Dalayest to daim No.						
Category °	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to daim No.						
Υ	SERAFIN, J.G., LIU, A.C., SEYEDMO	NIR,	1-20						
	S.R.: "Surface science and the silver-catalyzed epoxidation of e	thylene							
	an industrial perspective"								
	JOURNAL OF MOLECULAR CATALYSIS A:								
	CHEMICAL, vol. 131, 1998, pages 157-168, XP002230219								
	page 162, line 11 - line 39; figu	re 4							
Υ	EP 0 352 850 A (SHELL INT RESEARC	1-20							
	31 January 1990 (1990-01-31) claim 1								
A	CN 1 286 689 A (IDEMITSU KOSAN CO	1-20							
,	7 March 2001 (2001-03-07)	,							
	claim 1								
	_	/							
[] [her documents are listed in the continuation of box C.	Patent family members are listed	in anney						
		Y Patent family members are listed	iii dinox.						
		T later document published after the inte or priority date and not in conflict with	the application but						
A document defining the general state of the art which is not considered to be of particular relevance invention. *E* earlier document but published on or after the international invention.									
filing d		"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone							
which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the									
"O" document referring to an oral disclosure, use, exhibition or other means document is combined with one or more other such document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.									
	ent published prior to the international filing date but nan the priority date claimed	document member of the same patent family							
Date of the	actual completion of the international search	Date of mailing of the international sea	arch report						
6	February 2003	27/02/2003							
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l	Fax: (+31-70) 340-3016	Baston, E							

INTERNATIONAL SEARCH REPORT

Internatic pplication No PCT/US 02/37022

		PC1/US 02/3/022
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	WO 97 28142 A (ARCO CHEM TECH NL BV ;ARCO CHEM TECH (US)) 7 August 1997 (1997-08-07) claim 1	1-20
A	WO 97 10232 A (EASTMAN CHEM CO) 20 March 1997 (1997-03-20) claim 1	1-20

INTERNATIONAL SEARCH REPORT

nf tion on patent family members

Internatic pplication No
PCT/US 02/37022

				FC1/U3	02/3/022
Patent document cited in search report		Publication date		Patent family member(s)	Publication date
EP 0352850	, A	31-01-1990	AU	615561 B2	03-10-1991
	•		AU	3892089 A	25-01-1990
			BR	8903657 A	13-03-1990
			CA	1339317 A1	19-08-1997
			CN	1041591 A ,B	25-04-1990
			DE	68912440 D1	03-03-1994
			DE	68912440 T2	11-05-1994
			EP	0352850 A1	31-01-1990
			ES	2048273 T3	16-03-1994
			JP	2104579 A	17-04-1990
			JP	2779955 B2	23-07-1998
			KR	134882 B1	22-04-1998
			TR	24693 A	13-01-1992
CN 1286689	Α	07-03-2001	NONE		
WO 9728142	Α	07-08-1997	US	5625084 A	29-04-1997
			AT	202565 T	15-07-2001
			AU	721055 B2	22-06-2000
			AU	1545897 A	22-08-1997
			BR	9707465 A	20-07-1999
			CA	2244859 A1	07-08-1997
			CN	1210524 A	10-03-1999
			DE	69705391 D1	02-08-2001
			DE	69705391 T2	02-05-2002
			WO	9728142 A1	07-08-1997
			EP	0880513 A1	02-12-1998
			ES	2158486 T3	01-09-2001
			JP	11510817 T	21-09-1999
			· RU	2167872 C2	27-05-2001
			TW	448166 B	01-08-2001
			US	5686380 A	11-11-1997
WO 9710232	Α	20-03-1997	EP	0874837 A1	04-11-1998
			JP	2002510280 T	02-04-2002
			WO	9710232 A1	20-03-1997
			US	5908942 A	01-06-1999

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